Reply to Office Action of February 9, 2007

## AMENDMENTS TO THE CLAIMS

Docket No : 21058/0206453-US0

## Listing of Claims:

- 1 (Currently amended) A method comprising:
- a) providing one or more coded oligonucleotide probes, each coded oligonucleotide probe comprising an oligonucleotide attached to at least one unique nanocode wherein each nanocode comprises a detectable non-encoding feature, which detectable non-encoding feature comprises a feature tag pattern, and wherein the tag pattern provides a quality control check for detecting nanocodes and/or distinguishes target nucleotides from self-assembled coded oligonucleotide probe structures;
- b) contacting at least one target nucleic acid with the one or more coded oligonucleotide probes; and
- utilizing the feature tag to provide a quality control check for detecting nanocodes and/or distinguishes target nucleotides from self-assembled coded oligonucleotide probe structures; and
- identifying coded oligonucleotide probes that bind to the target nucleic acid using scanning probe microscopy (SPM) to detect the nanocode and the detectable non-encoding feature tag.
- 2. (Currently amended) The method of claim 1, wherein the one or more coded oligonucleotide probes comprise permutations of a linear order of nucleic acid residues, which linear order substantially represents all possible complementary sequences for a particular length of oligonucleotide.

Application No. 10/750,515

Amendment dated May 9, 2007

Docket No.: 21058/0206453-US0

Reply to Office Action of February 9, 2007

3. (Original) The method of claim 1, wherein the nanocode is selected from the

group consisting of carbon nanotubes, fullerenes, submicrometer metallic barcodes, nanoparticles

and quantum dots.

(Original) The method of claim 1, wherein the nucleic acid is attached to a

surface.

5. (Original) The method of claim 4, further comprising ligating adjacent coded

probes that are hybridized to the nucleic acid.

6. (Previously Presented) The method of claim 5, further comprising separating

ligated coded probes from the target nucleic acid and non-ligated coded probes.

(Original) The method of claim 6, wherein the ligated coded probes form

reading frames.

(Original) The method of claim 1, further comprising aligning the coded probes

on a surface by molecular combing.

9. (Previously Presented) The method of claim 1, wherein the scanning probe

microscopy is atomic force microscopy, scanning tunneling microscopy, lateral force microscopy,

Application No. 10/750,515 Docket No.: 21058/0206453-US0 Amendment dated May 9 2007

Reply to Office Action of February 9, 2007

chemical force microscopy, force modulation imaging, magnetic force microscopy, high frequency

magnetic force microscopy, magnetoresistive sensitivity mapping, electric force microscopy,

scanning capacitance microscopy, scanning spreading resistance microscopy, tunneling atomic force

microscopy or conductive atomic force microscopy.

10. (Previously Presented) The method of claim 2, further comprising determining

the nucleotide sequences of oligonucleotides that bind to the target nucleic acid.

11. (Previously Presented) The method of claim 10, further comprising

determining a nucleotide sequence of the target nucleic acid from the sequences of oligonucleotides

that bind to the target nucleic acid.

12. (Previously Presented) The method of claim 1, further comprising identifying

the target nucleic acid from the coded probes that bind to the target nucleic acid.

13. (Original) The method of claim 1, wherein two or more target nucleic acids are

present in a sample.

14. (Previously Presented) The method of claim 1, wherein at least two target

nucleic acids are contacted in the sample at the same time.

Application No. 10/750,515 Amendment dated May 9, 2007

Reply to Office Action of February 9, 2007

15. (Currently amended) The method of claim 1, wherein the detectable non-

Docket No.: 21058/0206453-US0

encoding feature tag is provided by a detectable feature tag associated with the nanocode.

16. (Currently amended) The method of claim 15 wherein the detectable non-encoding

feature tag comprises a start tag.

17. (Original) The method of claim 1, further comprising transforming the molecular

nanocode to form a decompressed nanocode.

18. (Currently amended) The method of claim 1, wherein the detectable feature tag is

comprises a checksum barcode segment.

19. (Currently amended) The method of claim 1, wherein the detectable feature tag

comprises a header segment and an encoding segment.

20. (Currently amended) A composition comprising at least one coded probe, each

coded probe comprising a probe molecule attached to at least one nanocode comprising a detectable

non-encoding feature; which detectable non-encoding feature comprises a feature tag pattern,

wherein the feature tag pattern provides has a property to provide a quality control check for

detecting nanocodes and/or distinguishes target nucleotides from self-assembled coded

oligonucleotide probe structures, the nanocode being detectable using a single molecule level

surface analysis method.

Reply to Office Action of February 9, 2007

21. (Previously Presented) The composition of claim 20, wherein the probe

molecule is an oligonucleotide, a polynucleotide, a nucleic acid, an antibody, an antibody fragment,

a genetically engineered antibody, a single chain antibody, a humanized antibody, a protein, a

receptor, a transcription factor, a peptide, a lectin, a substrate, an inhibitor, an activator, a ligand, a

hormone, a cytokine, a chemokine, or a pharmaceutical.

22. (Original) The composition of claim 20, wherein the probe molecule is an

oligonucleotide.

23. (Original) The composition of claim 20, wherein the nanocode is selected from

the group consisting of carbon nanotubes, fullerenes, submicrometer metallic barcodes.

nanoparticles and quantum dots.

(Currently amended) The composition of claim 20, wherein the detectable non-

encoding feature tag is comprises a start tag.

25. (Original) The composition of claim 20, wherein the nanocode is a compressed

nanocode.

26. (Original) The composition of claim 20, wherein the nanocode comprises

reading frames.

Application No. 10/750,515 Docket No.: 21058/0206453-US0 Amendment dated May 9, 2007

Reply to Office Action of February 9, 2007

27. (Original) The composition of claim 20, wherein the nanocode comprises a

header region and an encoding region.

28. (Original) The composition of claim 20, wherein the nanocode is detectable

using scanning probe microscopy (SPM).

(Currently amended) A system comprising:

a) a scanning probe microscope (SPM);

b) a surface: and

c) at least one coded oligonucleotide probe attached to the surface, wherein the coded

oligonucleotide probe comprises a nanocode comprising a detectable non-encoding feature, which

detectable non-encoding feature comprises a feature tag pattern, and wherein the feature tag pattern

provides has a property to provide a quality control check for detecting nanocodes and/or

distinguishes target nucleotides from self-assembled coded oligonucleotide probe structures, the

nanocode being detectable using SPM.

30. (Original) The system of claim 29, wherein the coded oligonucleotide probes

comprise ligated oligonucleotides.

31. (Original) The system of claim 30, wherein the ligated oligonucleotides form

reading frames.

Application No. 10/750,515

Amendment dated May 9, 2007

Docket No.: 21058/0206453-US0

Reply to Office Action of February 9, 2007

32. (Original) The system of claim 29, wherein the scanning probe microscope is an

atomic force microscope or a scanning tunneling microscope.

33. (Currently amended) The system of claim 29, wherein the detectable non-encoding

feature tag is comprises a start tag.

34. (Currently amended) The system of claim 29, wherein the nanocode is comprises a

compressed nanocode.

35. (Original) The system of claim 29, wherein the nanocode comprises reading

frames.

36. (Original) The system of claim 29, wherein the nanocode comprises a header

region and an encoding region.